

Extensive Plasmid Library to Prepare Tau Protein Variants and Study Their Functional Biochemistry

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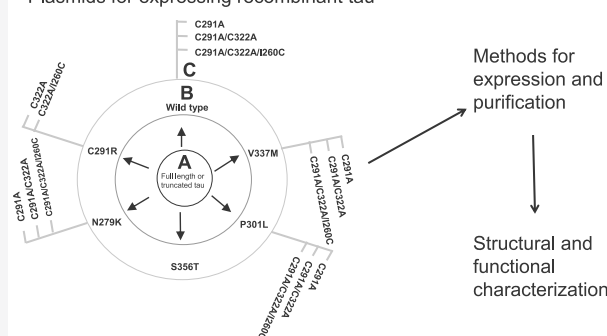


Supporting Information

ABSTRACT: Tau neurofibrillary tangles are key pathological features of Alzheimer's disease and other tauopathies. Recombinant protein technology is vital for studying the structure and function of tau in physiology and aggregation in pathophysiology. However, open-source and well-characterized plasmids for efficiently expressing and purifying different tau variants are lacking. We generated 44 sequence-verified plasmids including those encoding full length (FL) tau-441, its four-repeat microtubule-binding (K18) fragment, and their respective selected familial pathological variants (N279K, V337M, P301L, C291R, and S356T). Moreover, plasmids for expressing single (C291A), double (C291A/C322A), and triple (C291A/C322A/I260C) cysteine-modified variants were generated to study alterations in cysteine content and locations. Furthermore, protocols for producing representative tau forms were developed. We produced and characterized the aggregation behavior of the triple cysteine-modified tau-K18, often used in real-time cell internalization and aggregation studies because it can be fluorescently labeled on a cysteine outside the microtubule-binding core. Similar to the wild type (WT), triple cysteine-modified K18 formed paired helical filaments similar to those in Alzheimer's disease, sharing morphological phenotypes with WT tau-K18 filaments. Nonetheless, cysteine-modified tau-K18 filaments were significantly shorter ($p = 0.002$) and mostly wider than WT filaments, explainable by their different principal filament elongation pathways: vertical (end-to-end) and lateral growth for WT and cysteine-modified, respectively. Cysteine rearrangement may therefore induce filament polymorphism. Together, the plasmid library, the protein production methods, and the new insights into cysteine-dependent aggregation should facilitate further studies and the design of antiaggregation agents.

KEYWORDS: Tau, plasmid, MAPT mutations, Alzheimer's disease, frontotemporal dementia, expression and purification

Plasmids for expressing recombinant tau



INTRODUCTION

The microtubule (MT) associated protein tau (MAPT) gene, which encodes the tau protein, is located on chromosome 17q21.^{1,2} MAPT comprises 16 exons some of which are alternatively spliced to generate tau isoforms that are differentially expressed in tissues and in development.^{3,4} The adult human brain has six tau isoforms produced from the alternative splicing of exons 2, 3, and 10.^{1,5} Tau is comprised of a number of regions: the N-terminal projection domain, proline-rich domains in the midregion, and the C-terminal assembly domain, the latter containing the MT binding region [MTBR] and the C-terminus, separated by two proline-rich regions.^{1,5} The isoforms can be distinguished by the number of alternatively spliced inserts at the N-terminus (0, 1, or 2 N-terminus inserts) and in the MTBR (3 or 4 MTBR repeats).^{1,5} In normal adult humans, a 1:1 ratio of 4R to 3R is maintained.⁶ However, this isoform balance is disturbed in pathophysiological conditions, with tauopathies biochemically categorized

based on their tau isoform composition. Primary tauopathies (including corticobasal degeneration, progressive supranuclear palsy, and globular glial tauopathy) tend to have 4R tau isoforms (except Pick's disease with 3R tau), while secondary tauopathies such as Alzheimer's disease (AD) and chronic traumatic encephalopathy have a mix of 3R and 4R isoforms.^{7–11} The alternative splicing of exon 10 has been particularly linked to several tauopathies since it affects the ratio of three-repeat (3R) to four-repeat (4R) tau isoforms.¹²

The most-studied physiological function of tau is its support of axonal transport by binding to and modulating the stability

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of MTs.^{13,14} Tau binds to MTs at the α - and β -tubulin heterodimer interface using specific amino acid sequences, most of which are located within the MTBR.¹⁵ This generates a local hairpin structure involving the hexapeptide motifs ²⁷⁵VQIINK²⁸⁰ and ³⁰⁶VQIVYK³¹¹, with numbering corresponding to 441 amino acid-long full-length (FL) tau (Uniprot #P10636-8).¹⁵ Over 110 single nucleotide polymorphisms in *MAPT* have been reported, with more than a dozen associated with specific familial forms of frontotemporal dementia (FTD).^{16–19} Several of these disease-associated mutations can interfere with the protein's MT-binding function, by either potentiating or decreasing this activity.^{20–22} How specific *MAPT* mutations cause distinct effects on tau's MT interactions are not clearly understood, with further studies needed.

In disease conditions, tau can become hyperphosphorylated on specific residues leading to its detachment from MTs, thereby compromising axonal transport/assembly.²³ Following this loss of function, the soluble, unfolded phosphorylated tau proteins can self-polymerize to eventually form paired helical filaments (PHFs) and other filamentous aggregates enriched in β -sheet structures, via oligomer and protofibril intermediates.²⁴ These filaments are critical components of neurofibrillary tangles (NFTs), which are hallmarks of AD and other tauopathies.^{24,25} Tau aggregation can also occur through disulfide interactions involving the native cysteine residues, either at position 291 only (3R isoforms) or at 291 and 322 (4R isoforms).^{26–28} Intermolecular disulfide bonding is thought to promote aggregation, while intramolecular bonding inhibits aggregation.²⁹ 4R tau isoforms can therefore aggregate faster than 3R tau isoforms, possibly due to their capacity to form increasing polymers of disulfide-bonded aggregates relative to 3R tau isoforms whose disulfide-dependent aggregation is limited to dimerization.³⁰ However, the aggregation-prone ²⁷⁵VQIINK²⁸⁰ hexapeptide motif in 4R, but not 3R, tau isoforms may also contribute to the observed higher aggregation tendency of 4R tau, although little investigation has focused on this hypothesis.^{31,32}

Tau aggregation requires at least one of the hexapeptide motifs ²⁷⁵VQIINK²⁸⁰ and ³⁰⁶VQIVYK³¹¹ within the MTBR.^{31,33,34} Notably, a number of familial FTD-associated *MAPT* mutations, again located in the MTBR (e.g., P301L, V337M, N279K), are known to either accelerate or decrease tau aggregation kinetics and can form filaments of distinct structural phenotypes compared to the wild type (WT).^{20–22,35} Furthermore, some disease-associated mutations demonstrate stark differences in their ability to be “seeded” by filaments from the WT 4R tau MTBR (the K18 fragment), with P301L and P301S exhibiting robust template misfolding capabilities.^{35,36} This suggests that tau aggregation is conformation-dependent and that selected *MAPT* mutations may affect aggregation through distinct mechanisms. Moreover, the temporal and spatial progression of tau aggregate accumulation in AD and other tauopathies, which correlates well with cognitive decline, is thought to occur as a result of intercellular transfer and seeding of tau aggregates in cells.^{37,38} Specific *MAPT* mutations can influence this process through distinct effects on exogenous tau internalization and release.^{35,39} Moreover, tau aggregation can be a downstream event in AD (especially evident in familial AD due to mutations in the *APP* and *PSEN* genes), as well as in some other disorders (e.g., Niemann-Pick type C).⁴⁰ Together, these data demonstrate

that aggregation is a critical determinant of the pathophysiological functions of tau.

Much of the insight into the physiological and pathophysiological functions of tau (including MT binding, aggregation, and intercellular propagation) has been obtained using tau from recombinant sources, with many of these processes corroborated using human or animal tissues.^{22,23,41–45} These data establish recombinant technology as a validated strategy for generating functional tau proteins whose properties closely align with those of human brain tau protein. However, recombinant preparation of tau can be challenging and time-consuming, given that one needs to design and verify gene and plasmid sequences and perform polymerase chain reaction (PCR), restriction digestion, and ligation, before transforming a host of choice for expression. Even though some of these problems can be addressed by plasmid sharing between laboratories, the recipients often have to develop and optimize their own expression and purification protocols. These bottlenecks can unduly delay the preparation of recombinant tau. In an attempt to help address these shortcomings, we have generated a library of 44 sequence-verified plasmids for expressing FL human tau-441 (the longest tau isoform in the adult human brain) and truncated tau consisting of the 4R MTBR fragment (K18). These include plasmids for producing specific FTD variants of FL and K18 tau (V337M, N279K, P301L, C291R, and S356T), as well as cysteine-modified variants (C291A, C291A/C322A, C291A/C322A/I260C) of the FL and K18 fragment of WT and FTD tau variants.

Since FL 4R tau contains two native cysteine residues,²⁸ single and double cysteine modifications (usually replaced by alanine or serine) allow for studies of contributions of one or both native cysteine residues to the protein's properties of interest (MT binding, stabilization, and polymerization;^{46,47} filamentous aggregation and associated conformational changes^{26,28,48–52}). For studies aimed at investigating incipient tau aggregation and the intercellular propagation and toxicity of extracellular tau using real-time and/or kinetic assays, maleimide-based fluorophore labeling is, however, often used. This technique, first introduced by the Mandelkow groups,^{53,54} often involves triple cysteine modification where the native cysteine residues are removed (C291A/C322A) and a new cysteine residue introduced outside the central core of the MTBR (I260C) to minimize or avoid steric hindrance.^{35,49,53–58} Notwithstanding its increasing popularity, it has not been well studied if this triple cysteine modification affects tau aggregation kinetics, filament formation, and secondary structure. Importantly, answering this question will help provide a validated context to interpreting outcomes of experiments using these constructs. We therefore took advantage of the plasmid resources and the expression and purification methods developed to generate and carefully analyze the aggregation and Alzheimer-like filament formation paths and products of K18 C291A/C322A/I260C and K18 WT.

■ RESULTS AND DISCUSSION

Construction and Verification of Plasmids for Expressing FL and Truncated Tau and Their Respective Selected FTD Variants. pProEx-HTa plasmids encoding human WT FL or K18 tau (Figure S1) were created, and their sequences were verified by DNA sequencing. The coding sequences derived from the amino acid sequence for tau 1-441 (Uniprot #P10636-8) were generated by PCR from template

pCMV-tau plasmids and cloned into the *EheI* and *EcoRI* restriction sites of the pProEx-HTa plasmid as previously described.⁵⁹ DNA sequencing was performed externally by GATC Biotech AG, Konstanz, Germany. Protein expression was controlled by a *trc* promoter. The plasmid description is available in Table S1. Moreover, each plasmid had a cleavable N-terminal 6xHis tag for purification by immobilized metal affinity chromatography (IMAC; Figure S1). Each K18 plasmid also had a c-Myc tag inserted between the 6xHis tag and the genetic construct of interest, while the FL tau plasmids either had a C-terminus c-Myc tag or an N-terminus FLAG tag downstream of the 6xHis tag (Figure S1 and Table S1).

Specific FTD mutations can alter the physiological (including MT binding and stabilization and subcellular localization^{21,60–62}) and pathophysiological (e.g., phosphorylation, aggregation pathway, kinetics, and products^{21,22,35,63–66}) functions of tau. To increase the utility of our genetic resource library, the template WT plasmids were used to generate plasmids for expressing specific FTD variants of FL and K18 tau. We focused on selected FTD mutations spanning the MTBR, including some with relatively well characterized biochemical properties (e.g., V337M, N279K, and P301L^{21,36,65–69}) and others about which little is known (e.g., C291R and S356T^{70,71}). Successful generation of these constructs (Figure 1) was verified by DNA sequencing, with the codon changes and site-directed mutagenesis primers listed in Table S2.

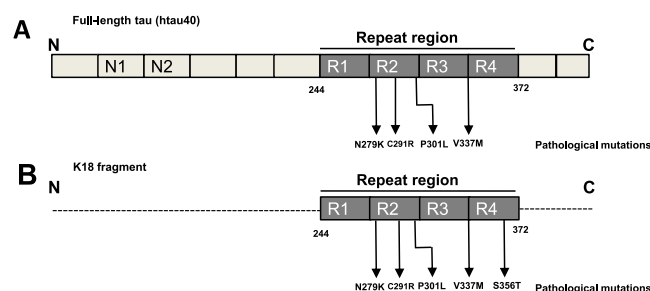


Figure 1. Schematic illustration of FL and truncated tau plasmid constructs carrying specific pathological *MAPT* mutations generated in this study. (A) In addition to the template plasmid for expressing WT FL tau-441, sequence-verified alternatives each carrying one of the FTD variants N279K, C291R, P301L, and V337M were constructed. The location of each pathological mutation in the MTBR is shown. (B) Plasmid constructs encoding the WT K18 fragment (the MTBR only) and its variants carrying one of the N279K, C291R, P301L, V337M, and S356T mutations created in this study. The location of each mutation is indicated.

Design and Verification of Plasmids for Expressing Cysteine Variants of FL, Truncated, and FTD Tau. The redox state of 4R tau, which has two native cysteine residues, can critically influence its properties.^{28,46,47,72} To support studies focused on understanding potential effects of the availability, quantity, and location of cysteine residues on tau functions, plasmids for expressing FL and truncated tau (both WT and FTD variants) bearing single (C291A), double (C291A/C322A), or triple (C291A/C322A/I260C) cysteine modifications were further generated (Figure 2 and Table S1). Codon changes made and primers used to generate these variants can be found in Table S3, with DNA sequencing primers in Table S4. Figure 3 shows a schematic illustration of the first (FTD mutations) and second order (cysteine

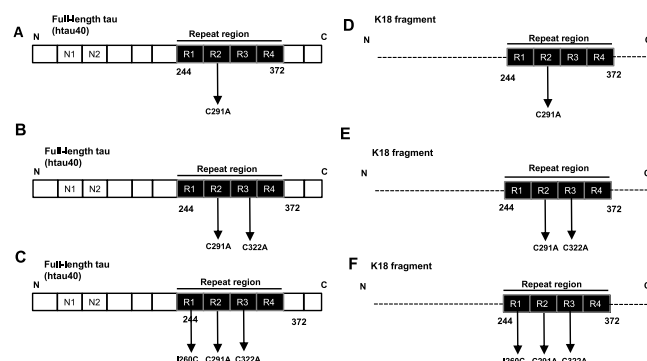


Figure 2. Schematic illustration of plasmid constructs created for the expression of WT and FTD versions of FL and K18 proteins containing specific cysteine modifications. Single (C291A), double (C291A/C322A), and triple (C291A/C322A/I260C) cysteine modifications in FL tau-441 (WT or FTD variants) are illustrated in (A), (B), and (C), respectively. (D), (E), and (F) describe the single, double, and triple cysteine modifications, respectively, in the K18 group of plasmids created in this study. The dashed lines denote the full length tau sequence which is removed in the truncated K18 construct.

modifications) plasmids generated based on each WT template construct.

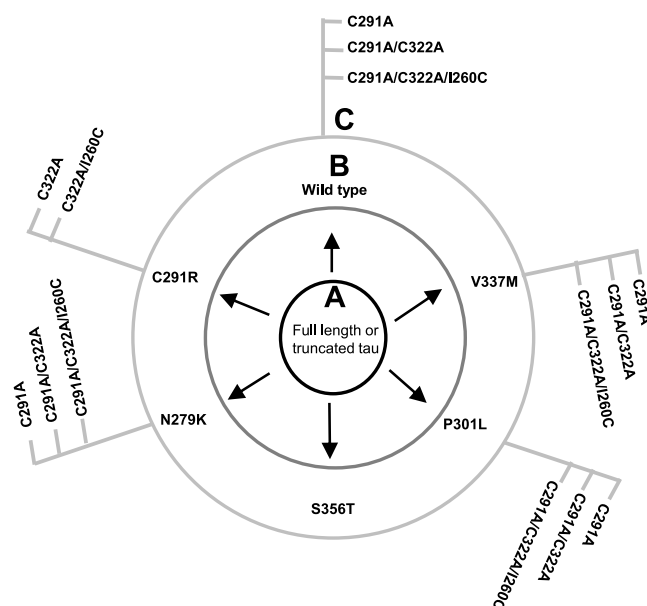


Figure 3. Incremental generation of plasmids expressing tau protein variants carrying specific FTD-associated mutations and cysteine modifications. Template plasmids for expressing WT FL and truncated tau were first created (A). Subsequently, mutagenesis of these base plasmids was used to generate variants encoding specific FTD mutations (B). Next, single, double, or triple cysteine modifications were introduced in the primary (WT) and secondary (FTD variants) plasmids to generate new forms of tau with variable cysteine content (C).

Expression and Purification of WT and Triple Cysteine-Modified FL Tau. To help simplify recombinant tau production, we developed a protocol for expressing and purifying selected constructs from our plasmid library. We adapted a high-yield method⁵⁹ to produce the WT and triple cysteine-modified forms of both FL tau and K18. Plasmids

transformed into BL21(DE3)*pRosetta *Escherichia coli* were expressed and purified by immobilized metal affinity chromatography (IMAC), taking advantage of the cleavable N-terminal 6xHis tags, as described (see the [Methods](#) section). Expressed protein variants were initially purified by IMAC. Tau-positive fractions, identified after SDS-PAGE and Western blotting, were further purified by size exclusion chromatography (SEC), and the purest fractions were selected ([Figure 4](#)).

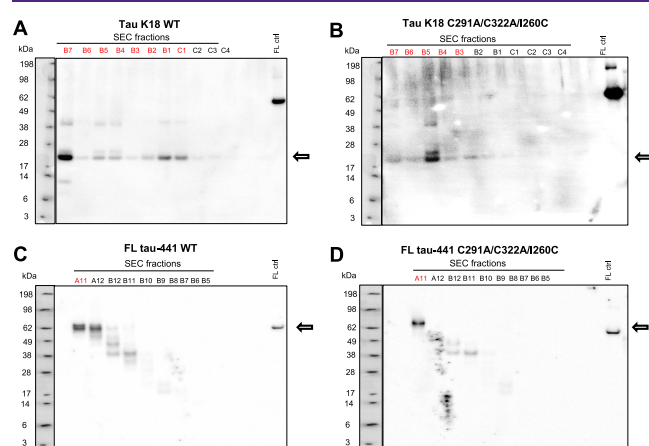


Figure 4. Expression and purification of WT FL and truncated tau K18 and their triple cysteine modified variants. Representative characterization of tau purification by Western blotting. K18 WT (A) and K18 triple cysteine-modified (B) were analyzed by Western blotting using the 77G7 antitau antibody (epitope = aa 316–355), while the WT (C) and triple cysteine modified (D) FL tau were probed with Tau12 (epitope = aa 6–18). A previous preparation of WT recombinant FL tau-441 was used as a control on each blot. The alphanumeric labeling of lanes refers to the analyzed fractions from size exclusion chromatography that were analyzed by Western blotting. Bands corresponding to tau which is in the conformation of interest are indicated by black arrows. Fractions labeled in red were the purest/highest yield selected for further processing. [Figures S2 and S3](#) show Western blot characterization of the initial IMAC-based purification. Size characterization of purified tau forms using SEC can be found in [Figure S4](#).

Cleavage of the 6xHis tag was achieved by overnight dialysis in the presence of tobacco etch virus (TEV) protease, followed by a second IMAC purification resulting in the 6xHis tag bound to the column, liberating the cleaved protein as either tau-K18 with a c-Myc tag or FL tau-441 with a FLAG tag.⁵⁹

The IMAC-purified 6xHis-tagged tau K18 fragments migrated on SDS-PAGE gels with electrophoretic mobility corresponding to ~22 kDa and were recognized by the 77G7 ([Figure 4A,B](#)) and K9JA⁵⁹ antitau antibodies both selective for the MTBR (epitopes = amino acids (aa) 316–355 for 77G7 and 244–372 for K9JA). Cleavage of the 6xHis tag was demonstrated by lost reactivity to the anti-His antibody,⁵⁹ reducing the apparent molecular weight of the K18 peptides to ~17 kDa by Western blotting ([Figure S2](#)). WT and triple cysteine-modified FL tau were also purified to near homogeneity with apparent sizes of ~62 kDa on SDS-PAGE/Western blot, with strong reactivity to the N-terminus antitau antibody Tau12 (epitope = aa 6–18; [Figure 4C,D](#)) and the MTBR-binding K9JA ([Figure S3](#)). For characterization of purified tau species using SEC, see [Figure S4](#).

Thioflavin T Assay Shows That Triple Cysteine Modification Decreases the Aggregation Propensity of WT Tau K18. A ThT assay was developed to study if the

triple cysteine modification affected the aggregation kinetics of K18 WT. K18 WT had an initial mean signal of ~1200 relative fluorescent units (RFU), which after a lag phase of ~1.5 h, increased sharply until reaching a plateau at ~11,400 RFU just after 5 h ([Figure 5A,B](#)). On the contrary, K18 C291A/C322A/I260C

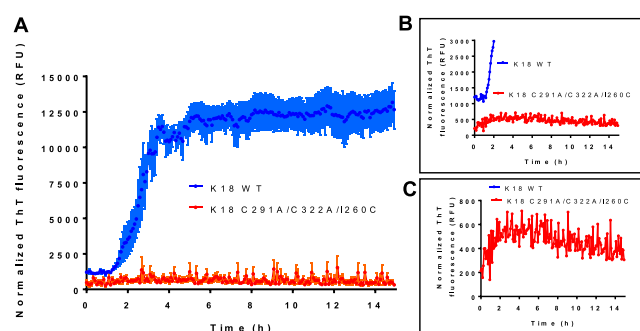


Figure 5. Aggregation kinetics of WT and triple cysteine-modified tau K18 in the ThT fluorescence assay. (A) Aggregation kinetics of the tau K18 variants each normalized to corresponding signals from the constituent buffer only + ThT (negative control). Normalization was achieved by subtracting the ThT signal generated by buffer only at each time point from the signals generated by the tau K18 constructs. Data are expressed as mean \pm SEM, $n = 4$ and $n = 11$ for K18 WT and K18 C291A/C322A/I260C, respectively. (B) Magnified image focusing on values ≤ 3000 RFU to highlight the shape of the kinetic curve for K18 C291A/C322A/I260C. (C) ThT fluorescence values ≤ 800 RFU demonstrate that K18 C291A/C322A/I260C recorded a brief rise in aggregation kinetics which then plateaued.

I260C had severely reduced ThT kinetics: an initial mean signal at ~200 RFU increased slowly until reaching a peak at ~600 RFU after 3 h ([Figure 5A–C](#)). The kinetic behavior of K18 C291A/C322A/I260C was more visible by focusing on RFU values < 800 , which showed that the peptide had an initial small increase in ThT fluorescence which immediately stabilized after 3 h: no discernible lag phase was recorded ([Figure 5C](#)). The data suggest that the triple cysteine modification decreases the aggregation rate of tau K18 ~20 fold in nonreducing conditions.

WT Tau K18 Forms Filaments Faster than Its Triple Cysteine-Modified Variant. Co-incubation of amyloidogenic proteins with the ThT dye can lead to unintended effects on aggregation.⁷³ To rule out the possibility that the decreased aggregation kinetics of K18 C291A/C322A/I260C was due to anomalous interactions with ThT, a dye-free aggregation assay was performed. No filament formation was recorded for either sample incubated for 15 h at 37 °C without shaking. Consequently, the incubation time was extended to promote filament assembly. After 5 days in the same conditions, K18 WT, but not K18 C291A/C322A/I260C, formed filamentous aggregates ([Figure 6A,B](#)). This corroborates the ThT-monitored aggregation data in [Figure 5](#) that the triple cysteine modification decreases the aggregation kinetics of K18 WT in nonreducing conditions.

End Point Tau K18 Aggregates Have Converging and Diverging Structural Phenotypes. Incubation time for aggregation reactions was further increased to 16 days to enable both tau variants to form mature filamentous aggregates. TEM analysis showed that K18 WT and K18 C291A/C322A/I260C formed filaments with both converging and diverging phenotypes. Both variants formed paired helical filaments (PHFs) with varying degrees of twist ([Figure 7A–](#)

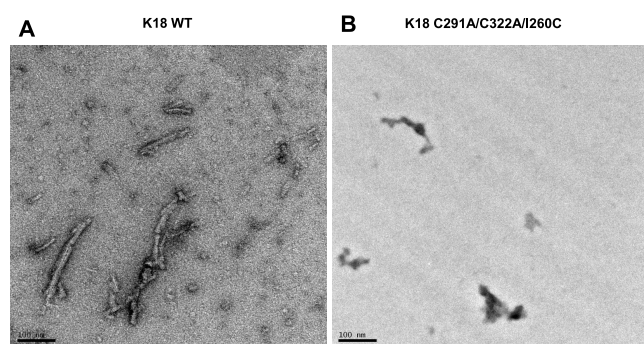


Figure 6. Structural characterization of tau K18 aggregation products in 5-day aggregation reactions. Representative TEM micrographs for tau K18 WT (A) and K18 C291A/C322A/I260C (B). Images were taken using identical TEM settings at 25000 \times magnification. Scale bars = 100 nm.

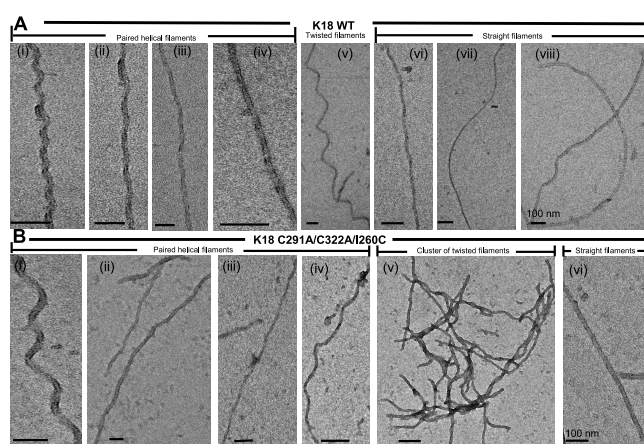


Figure 7. Characterization of the aggregation products of tau K18 variants after 16 days of incubation. Representative TEM micrographs for tau K18 WT (A) and K18 C291A/C322A/I260C (B) are shown, demonstrating both shared and distinct morphological features between filaments formed by the two constructs. Images were taken using identical TEM settings at 12000 \times magnification. Scale bars = 100 nm for each panel.

iv,Bi–iv). In addition to classical PHFs (Figure 7Aiii,iv), K18 WT formed multistriated ribbons (multiple PHFs interwoven together; Figure 8Ai) and ribbons with central depression (multistriated ribbons of uneven widths; Figure 7Aii), consistent with previous observations.⁷⁴ Furthermore, a few twisted filaments lacking the classical PHF arrangement were observed (Figure 7Av). Moreover, in K18 WT preparations, two types of straight (nontwisted) filaments (SFs) were recorded: those with the regular linear appearance (Figure 7Avi) and others with a curved phenotype (Figure 7Avii,viii). K18 C291A/C322A/I260C formed mostly PHFs which often made lateral contacts with others and sometimes formed clusters (Figure 7Bi–v). A few SFs were also observed (Figure 7Bvi).

WT tau K18 filaments were significantly longer than those of K18 C291A/C322A/I260C ($p = 0.002$, two-tailed Mann–Whitney test; mean lengths for WT and C291A/C322A/I260C = 1457.00 ± 76.33 nm and 1069.00 ± 86.73 nm respectively; Figure 8A). On the contrary, several K18 C291A/C322A/I260C filaments appeared to be wider than those of the WT (mean width for WT and cysteine-modified = 16.80 ± 1.27 nm and 20.38 ± 1.95 nm) although this difference was

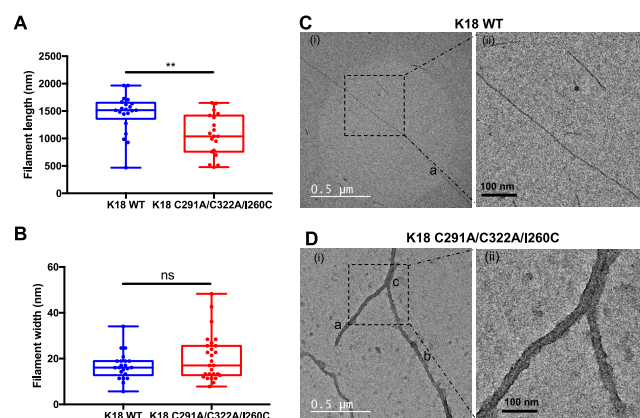


Figure 8. Quantitative characterization of K18 WT and K18 C291A/C322A/I260C filaments. (A) Box plot of filament lengths for the two tau K18 variants. Two-tailed Mann–Whitney test, $** = p = 0.002$. Mean lengths for WT and cysteine-modified K18 filaments = 1457.00 ± 76.33 nm and 1069.00 ± 86.73 nm, respectively. Median lengths for K18 WT and K18 C291A/C322A/I260C = 1515.00 and 1038.00 nm, respectively. The filament length analysis included $n = 21$ data points for K18 WT and $n = 19$ for K18 C291A/C322A/I260C. (B) Box plot of filament widths for the tau K18 variants. Two-tailed Mann–Whitney test, $p = 0.376$; ns = not significant. Mean widths for K18 WT and K18 C291A/C322A/I260C = 16.80 ± 1.27 nm and 20.38 ± 1.95 nm, respectively. Median widths = 16.07 and 17.05 nm for K18 WT and K18 C291A/C322A/I260C, respectively. The filament width analysis included $n = 22$ data points for K18 WT and $n = 28$ for K18 C291A/C322A/I260C. (C) (i) An example TEM image showing the long and slender appearance of the K18 WT filament. The image shown here has a length of 1966.59 nm and a width of 5.68 nm. Position where the width measurement was taken is marked “a”. (ii) A digitally zoomed image highlighting the long and thin appearance of the filament structure shown in (i). (D) (i) A representative TEM micrograph demonstrating lateral connections between two K18 C291A/C322A/I260C filaments. The upper sections of these filaments had coalesced to form a thicker filament. The filament on the left is 1038.08 nm long and 36.16 nm wide (position marked “a”), while the one on the right is 1519.06 nm long and 22.72 nm wide (position marked “b”). The width at the point of coalescence (marked “c”) is 48.30 nm. (ii) Digital zoom showing a magnified view of the lateral interactions observed in (i). The images in (C)(i) and (D)(i) were both acquired using identical TEM settings on the same day at 12000 \times magnification.

not found to be significant ($p = 0.376$, two-tailed Mann–Whitney test; Figure 8B). Careful observation of TEM images suggested that contrary to filament growth for K18 WT, which appeared to occur mainly through a vertical (end-to-end) growth mechanism (due to their generally long and slender morphology), K18 C291A/C322A/I260C had a lateral growth mechanism evidenced by regular lateral contacts between filaments, similar to those reported for amyloid β with an N-terminal cysteine.⁷⁵ Quantitatively, 90% (45/50) of fibrils formed by K18 C291A/C322A/I260C made lateral contacts with at least one other fibril. To the contrary, only 12% (5/41) lateral contacts were recorded for K18 WT. Figure 8C shows a typical TEM field of view for K18 WT filaments, demonstrating their long appearance without lateral contacts. Figure 8D shows an example of laterally interacting K18 C291A/C322A/I260C filament: wider but shorter than the K18 WT filament in Figure 8C. Our finding of vertical growth mechanism for K18 WT filaments is supported by a recent study that demonstrated this mechanism for FL tau-441.⁷⁶

Aggregation of Triple Cysteine-Modified Tau K18 Is Characterized by Increased β -Sheet Structures. It is well-known that the aggregation of K18 WT results in secondary structure rearrangement toward ordered β -sheet conformers.^{53,59,77,78} We therefore investigated if K18 C291A/C322A/I260C also had this property. K18 C291A/C322A/I260C monomers had negative CD spectra with the mean minimum peak at 199.00 ± 0.55 nm, similar to that of K18 WT at 200.70 ± 0.15 nm (Figure S5A,B), suggesting predominant random coil conformations^{59,77,78} and indicating that the triple cysteine modification does not alter the unfolded structure of tau K18 monomers. Following heparin-induced aggregation without shaking, the K18 C291A/C322A/I260C spectra shifted toward longer wavelengths, with new mean peaks of 202.90 ± 1.07 nm at day 4 and 206.90 ± 0.03 nm at day 8 (Figure 9A and Table

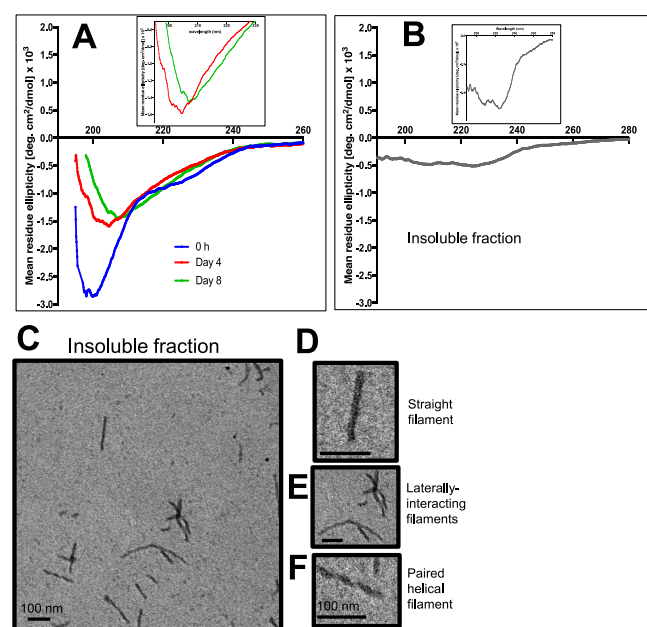


Figure 9. Aggregation of tau K18 C291A/C322A/I260C is characterized by progressive β -sheet enrichment. (A) CD analysis of time-dependent secondary structure changes in tau K18 C291A/C322A/I260C during aggregation. Mean negative peak signals increased from 196.60 ± 0.71 nm at day 0 to 202.90 ± 1.07 nm and 206.90 ± 0.03 nm at days 4 and 8, respectively. The inset image highlights peak intensity transition from day 4 to day 8. Each plot shows the mean of $n = 3$ repeats with quantitative data given in Table S5. (B) Mean CD spectrum of secondary structure content of the insoluble fraction of aggregated tau K18 C291A/C322A/I260C showing peak intensity at 224.00 ± 1.10 nm ($n = 2$, quantitative data provided in Table S6). The inset highlights the spectrum peak on a shorter vertical axis. (C) TEM characterization of the insoluble aggregate fraction from (B). Similar to data shown in Figures 8 and 9, K18 C291A/C322A/I260C formed a mix of PHFs and SFs. The images in (D), (E), and (F) are zoomed in regions from (C). Scale bars = 100 nm.

S5). Insoluble aggregates (resuspended pellets) had negative spectra with a mean peak at 224.00 ± 1.10 nm (Figure 9B and Table S6). The K18 C291A/C322A/I260C insoluble aggregate fraction consisted of abundant filaments (Figure 9C), including SFs (Figure 9D) and PHFs (Figure 9F) and laterally interacting filaments (Figure 9E) similar to those described in Figures 7 and 8. Together, β -sheet content in K18

C291A/C322A/I260C increased progressively with aggregation.

CONCLUSION

The preparation of recombinant forms of tau requires sequence-verified plasmids for overexpressing the protein in suitable expression hosts such as *E. coli*.^{59,79,80} The main aim of this study was to design, produce, and characterize a large library of plasmids to simplify the preparation of recombinant FL and truncated tau protein variants. A total of 44 sequence-verified plasmids were generated. These include plasmids for expressing WT and FTD tau (V337M, N279K, P301L, C291R, S356T) and their respective cysteine-modified variants (C291A, C291A/C322A, C291A/C322A/I260C) on FL and K18 backgrounds. Moreover, we have established protocols that ensure high-level expression and also demonstrate the purification and initial biochemical characterization of a representative selection of FL and truncated tau genetic constructs.

In this study, we focused on FL tau-441 and K18, both of the 4R tau background. More than 110 MAPT mutations have been described, with more of such mutations occurring in 4R than 3R tau isoforms.¹⁹ 4R tau has superior MT binding ability compared with 3R tau¹⁴ and is a common constituent of PHFs and NFTs isolated from the brains of individuals who died with many forms of neurodegenerative tauopathies.^{67,81–83} Moreover, truncation of 4R tau into fragments containing whole or part of the MTBR is a post-translational modification with probable biomarker importance.^{84,85} 4R tau therefore offers an opportunity to study the molecular origins of MT binding, aggregation, and filament formation, with direct relevance to the protein's physiological and pathophysiological functions.

Verified Plasmids and Protocols for Expressing and Purifying FL and Truncated Tau. Our comprehensive plasmid library is meant to help address common bottlenecks in plasmid sharing by providing ready-to-use genetic resources with accompanying protocols for generating FL and truncated tau variants. With the described protocols, the expression and purification process is achievable in 2 or 3 days for 6xHis-tagged and 6xHis tag-free tau, respectively, far shorter than other published protocols.^{79,80} We expect that this first-of-a-kind open access genetic resource library and the accompanying methods will facilitate further studies. The expression and purification protocols described are representative for the FL and truncated tau forms created, as demonstrated above for the cysteine-modified constructs and previously for the N279K, V337M, and C291R variants.^{35,58,59,86}

Impact of Triple Cysteine Modification on Tau K18 Aggregation and Filament Formation. Commonly used fluorescent protein tags such as green fluorescent protein can lead to unintended cellular effects,^{87,88} which in addition to their large molecular weights (~ 27 kDa) makes them unideal as reporter systems for internalization and release of exogenous tau. For this reason, the triple cysteine-modified forms of tau have become an increasingly popular set of tools for studying the aggregation, cellular internalization, release, and propagation of these proteins in real time because of the advantage of labeling a single cysteine residue located outside the MTBR with fluorescent compounds.^{49,53–57} In addition, we recently reported that specific labeling of triple cysteine-modified tau K18 with the maleimide derivatives *N*-ethyl maleimide or Alexa Fluor C3-maleimide in cold conditions generates stable

tau oligomers suitable for biochemical, cellular, and electrophysiological studies.^{35,58,89,90} However, it is unknown how the *in vitro* aggregation mechanism compares to that of K18 WT. We therefore took advantage of the resources and methods created to investigate potential effects of the triple cysteine modification on the aggregation paths and products of tau K18. Some earlier studies reported that 4R tau isoforms have decreased aggregation kinetics compared to 3R isoforms, which may be due to the two native cysteine residues in 4R tau forming aggregation-incompetent intramolecular disulfide bonds (the so-called *compact monomer* conformation).²⁹ However, others have argued that intramolecular disulfide bonds may not have significant effects on 4R tau aggregation and filament formation.^{49,52} Here, we sought to help address this unresolved question by studying if the number of cysteine residues influences tau K18 aggregation kinetics and filament structure. While some previous studies have compared 3R and 4R tau isoforms and constructs directly,²⁹ the deletion of an entire repeat domain may have affected their experimental outcomes, for instance, given the crucial pro-aggregation role of the ²⁷⁵VQIINK²⁸⁰ hexapeptide motif in the second repeat domain which is absent in 3R tau.^{31,91} Given that the K18 region is required for both disulfide-dependent and disulfide-independent tau aggregation,⁹² we studied the double-cysteine WT fragment and its C291A/C322A/I260C single-cysteine variant which differ in the number of cysteine residues. Although we cannot rule out potential effects of cysteine positional changes in this variant, the present study was focused on understanding disulfide bonding contributions to tau K18 aggregation based on the number of cysteine residues. This enabled detailed biochemical characterization of the *in vitro* aggregation properties of K18 C291A/C322A/I260C. Our findings indicate that tau K18 WT, with two cysteine residues, aggregates into filaments more rapidly than its single-cysteine variant. This is in agreement with a previous study which demonstrated that the 3R FL tau (which has a single native cysteine) forms less abundant dimers and higher order oligomers compared to the 4R FL tau.⁹² An explanation for this result is that as K18 C291A/C322A/I260C contains a single cysteine, it can only form disulfide-dependent dimers without disulfide-dependent higher-order aggregates compared to the WT which can aggregate into larger structures via the disulfide-dependent pathway²⁷ (Figure 10). K18 WT therefore had a faster aggregation due to combined effects of disulfide-dependent and -independent aggregation processes compared to K18 C291A/C322A/I260C with weaker disulfide-dependent activity.

An alternative interpretation for our results is that the ability of tau K18 to form filamentous aggregates is independent of whether a cysteine residue is located within the first, second, or third repeat domains. Previous studies on the contribution of disulfide bonds to 4R tau aggregation modified single or double cysteine residues to nonpolar amino acids like alanine.^{26–28,51} Nonetheless, effects of introducing cysteine residues outside the second and third repeat domains have hitherto been unknown. We have shown here that restricting cysteine residues to the second and third repeat regions (for K18 WT) is not a requirement for Alzheimer-like filament formation *in vitro*. Rather, aggregation characterized by β -sheet-dependent filament assembly is attainable, although at a reduced rate, when a single cysteine residue is placed in the first repeat domain.

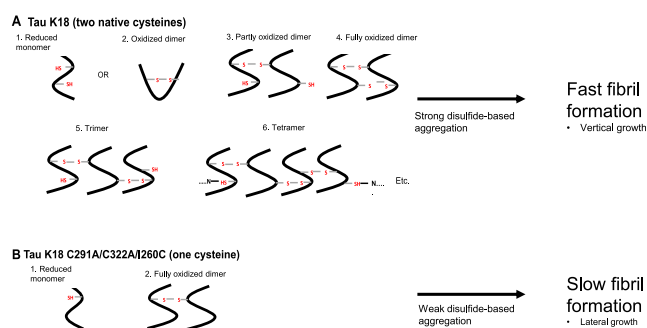


Figure 10. A proposed model of cysteine-dependent tau K18 aggregation in nonreducing conditions. (A) Monomeric tau K18 WT, with its two native cysteine residues, can undergo either intramolecular disulfide bonding to form “compact monomers” or intermolecular disulfide bonding to form aggregation-competent dimers. Partially oxidized dimers, but not fully oxidized dimers, can undergo further cysteine-dependent aggregation with reduced monomers to form trimers, tetramers, and correspond larger aggregates. (B) Tau K18 C291A/C322A/I260C contains a single cysteine residue and for that matter can only form an oxidized dimer, which cannot further aggregate via the cysteine-dependent pathway. Since both proteins have intact hexapeptide motifs that also modulate aggregation, it is proposed that the strong cysteine-dependent aggregation tendency of tau K18 explains its overall higher aggregation and filament-forming tendency compared to K18 C291A/C322A/I260C.

An important finding from this study is that a specific rearrangement of native cysteine residues resulting in a single cysteine outside the MTBR can drastically decrease the aggregation kinetics and delay filament assembly in tau K18. Similarly, modification of the C291 residue of K18 WT to serine inhibited the peptide’s oligomerization competence.²⁷ The strategy of modifying cysteine residues may therefore inform the design of effective approaches to inhibit 4R tau aggregation. Compared to the computer-guided random amino acid shuffling approach used to generate aggregation-compromised scrambled proteinopathies,^{93,94} cysteine modification is a more reproducible strategy that uses a nonpharmacological approach to inhibit tau K18 aggregation, with its effectiveness independent of external agents. Given the physiological and pathophysiological importance of 4R tau,^{16,95,96} a viable approach to block its disease-driving aggregation will be useful for targeting most tauopathies.

The observed similarities and differences in filament morphology for K18 WT and K18 C291A/C322A/I260C suggest that the two protein variants have both shared and divergent conformational changes leading to filamentous aggregation. The mechanisms underlying these similar-yet-diverse properties are not currently understood. However, a few features stand out. K18 WT mostly formed long and thin filaments. Filament assembly by this peptide involved vertical association between two short filaments, as also reported in a different study.⁷⁶ In contrast, K18 C291A/C322A/I260C filament elongation was dominated by lateral interactions of filaments. As demonstrated by TEM imaging (Figure 8D), coalescence of laterally associated filaments appeared responsible for the generally wide nature of the resulting mature filaments and may explain the frequently short appearance of K18 C291A/C322A/I260C filaments. Importantly, the lateral growth mechanism has been described for amyloid β with an N-terminal cysteine,⁷⁵ which highlights the importance of a single cysteine residue for this aggregation pathway. To our

knowledge, however, this is the first description of this mechanism for a tau form. Notwithstanding their apparent distinct principal filament formation mechanisms, K18 WT and K18 C291A/C322A/I260C formed some filaments with the same structural phenotypes. This raises the possibility that filaments of the same structural phenotypes can be formed through apparent heterogeneous aggregation paths.

Based on our findings, we propose a new model of cysteine-dependent tau aggregation (Figure 10). In nonreducing conditions, K18 WT can form either an intramolecular disulfide bond (compact monomer) or an intermolecular disulfide-based dimer as its least complex aggregate. Compact monomers do not appear to impede aggregation, potentially due to a low abundance or unfavorable formation in the presence of heparin. Dimers can associate with reduced monomers to form trimers and corresponding higher aggregates by disulfide linkages. Partially oxidized dimers/trimers/tetramers/larger oligomers can also cointeract through disulfide bridges, therefore accelerating the rate of aggregation (Figure 10). For K18 C291A/C322A/I260C, compact monomer formation is not feasible, but a dimer is the only cysteine-dependent aggregate it can form (Figure 10). Since both constructs can theoretically undergo cysteine-independent aggregation to the same extent (each has intact aggregation-promoting hexapeptide motifs in the MTBR), we hypothesize that the stronger aggregation tendency of K18 WT is conferred by its higher potential of cysteine-dependent polymerization. It is currently unclear if these disulfide linkages have any association with the filament assembly mechanisms observed.

A limitation of this work is that even though the K18 fragment allows for accelerated aggregation, using this truncated construct inevitably simplifies the otherwise more complicated process for FL tau. Therefore, not all aspects of the functional characterization described here for tau K18 may apply to FL tau. Nonetheless, the described plasmids and biochemical protocols will allow complementary FL tau versions of the K18 constructs to be produced and characterized.

In conclusion, we expect that the tau plasmid library and the associated expression and purification protocols described herein will facilitate future studies into the structural and molecular determinants of the protein's MT binding in physiological conditions and aggregation in multiple diseases and aid the design of effective inhibition agents. Moreover, the new insights provided into the mechanistic basis of tau K18 filament polymorphisms should pave the way for further studies exploring this knowledge for therapeutic gains.

METHODS

Materials. Heparin (#H3393), sodium phosphate dibasic (#S9763), sodium phosphate monobasic dihydrate (#4269), and thioflavin T (#T3516) were obtained from Sigma-Aldrich (St. Louis, Missouri, USA). The QIAprep Spin Miniprep kit and the Q5 site-directed mutagenesis kit (#E0554S) were bought from QIAGEN GmbH (Hilden, Germany) and New England Biolabs (Ipswich, Massachusetts, USA), respectively. The Bicinchoninic acid assay kit (#786-570) was obtained from G-Biosciences (St. Louis, Missouri, USA). The K9JA total tau antibody (#A0024) was purchased from Dako (now Agilent), 77G7 (#816701) and Tau12 (#806501) were purchased from BioLegend (San Diego, California, USA), and fast flow resin with chelating sepharose (#17-0575-01) was purchased from GE Healthcare (Buckinghamshire, UK). NuPAGE 4–12% Novex Bis-Tris gels (#NP0321), 4× LDS (lithium dodecyl sulfate)

buffer (#NP0007), 20× MES (2-(*N*-morpholino)ethanesulfonic acid) SDS running buffer (#NP0002), and 10× sample reducing agent (#NP0004) were purchased from ThermoFisher. HRP-labeled secondary antibody (#W4021) was purchased from Promega. Isopropyl β -D-1-thiogalactopyranoside (IPTG; #16758) and DNase I (#04716728001) were purchased from Sigma, while BugBuster (#70921) and protease inhibitor cocktail (#11836153001) were purchased from Merck. BL21(DE3)*pRosetta cells and TEV protease were kind gifts of Prof. David Roper and Dr. Deborah Brotherton, respectively, both at the University of Warwick, Coventry, UK.

Plasmid Cloning, Protein Expression, and Purification.

Plasmid design, cloning, site-directed mutagenesis, transformation, and DNA sequencing have been described elsewhere.⁵⁹ Amino acid changes made and primers used for site-directed mutagenesis have been listed in Tables S2 and S3. Plasmids meant for open-access sharing were prepared by transformation into BL21(DE3)*pRosetta cells and amplified by overnight growth in Luria–Bertani broth containing ampicillin (100 μ g/mL) and chloramphenicol (35 μ g/mL), and the DNA was isolated using the QIAprep Spin Miniprep kit following the manufacturer's instructions. DNA concentration was determined on a NanoDrop NP60 spectrophotometer. Mutagenic changes were confirmed by aligning DNA sequencing chromatograms against the template plasmid sequences using Genome Compiler (Twist Bioscience, California, USA). Plasmid DNA aliquots are available for sharing, allowing their transformation into the host of choice. Details of plasmids can be found in Table S1.

Plasmids transformed into BL21(DE3)*pRosetta cells were expressed and first purified by IMAC as described.⁵⁹ Briefly, an overnight culture of 10 mL of selective LB broth inoculated with single colonies of BL21(DE3)*pRosetta cells encoding the tau plasmid of interest was used to inoculate a 1 L fresh selective LB broth. The cultures were grown at 37 °C with 180 rpm shaking until the OD₆₀₀ reached 0.6–0.7. After inducing expression with 0.5 mM IPTG, the cultures were returned to the 37 °C shaking incubator for a further 1 h. The cell lysates were harvested by centrifuging the cultures at 9800 \times g for 10 min at 4 °C. The recovered pellets were resuspended in 50 mM sodium phosphate buffer pH 7.5 and frozen at –20 °C until use. Immediately before purification, the lysates were heated at 70 °C in the presence of the following reagents added per 50 mL lysate: 5 mL of 1× BugBuster, 10 mg DNase I, and 1 tablet protease inhibitor cocktail. The mixtures were kept at room temperature for 1 h and then sonicated (70% power, 1 min) in a Bandelin Sonopuls 2070 sonicator and centrifuged (48000 \times g, 30 min, 4 °C). The supernatant was decanted and filtered (0.2 μ m) before IMAC purification.

Chelating Sepharose resin packed into a Ni-NTA column (Bio-Rad) was charged with 10 mM NiCl₂/CH₃COONa pH 4.0 buffer and equilibrated with two column volumes (CVs) of buffer A (50 mM sodium phosphate buffer pH 7.0, 500 mM NaCl, 10 mM imidazole). After applying the crude extract (input), unbound protein was removed in a two-step washing protocol, first with two CVs of buffer A and then 2 CVs of buffer B (50 mM sodium phosphate buffer pH 7.0, 500 mM NaCl, 25 mM imidazole). His-tagged proteins were eluted with buffer C (50 mM sodium phosphate buffer pH 7.0, 500 mM NaCl, 500 mM imidazole, 10 mM dithiothreitol). Selected fractions, as analyzed by SDS-PAGE and Western blotting, were pooled and concentrated by ultrafiltration using 3 and 10 kDa cutoff centrifugal filter devices (Amicon) for K18 and FL tau, respectively. All constructs were initially further purified by size exclusion chromatography (SEC) on a Superdex S200 10/300 GL column (#17-5175-01, GE Healthcare) running on an Ethan LC system (GE Healthcare) with 1× PBS pH 7.4 as the running buffer. The molecular weights of the fractions of interest were estimated using the elution volumes of the following size markers from Sigma (#MWGF70-1KT): blue dextran (2000 kDa; void volume), bovine serum albumin (66 kDa), carbonic anhydrase (29 kDa), cytochrome C (12.4 kDa), and aprotinin (6.5 kDa). The optimized process showed that a SEC polishing step was not necessary for K18 (data not shown). For all constructs, the 6×-His tag was removed by overnight dialysis against dialysis buffer (50 mM Tris HCl pH 7.5, 100 mM NaCl) in the

presence of 25 $\mu\text{g/mL}$ TEV protease. The cleavage products were separated by applying the dialyzed preparation to a charged resin equilibrated as previously and collecting the cleaved tau as the flow-through.

Western Blot. Equal volumes of SEC fractions were diluted in the LDS sample buffer and the sample reducing agent and run in 4–12% Bis-Tris protein gels in MES SDS running buffer (gel and buffers were purchased from ThermoFisher Scientific; see the [Materials](#) section above). The gels were subsequently equilibrated in transfer buffer (20% methanol) for 25 min. Proteins were transferred to a polyvinylidene fluoride membrane which had been previously activated in pure methanol for 30 s and soaked in the transfer buffer for another 25 min. Transfer was performed in a semidry blot instrument at 0.8 mA/cm² for 60 min at room temperature. The membrane was then blocked for 1 h at room temperature using PBST-BSA (PBS with 0.05% Tween 20 and 1% BSA) with gentle shaking. Primary antibody incubation was performed overnight at 4 °C at a ratio of 1 μg of antibody per 1 mL of PBST-BSA. Afterward, the membrane was washed three times with PBST (PBS with 0.05% Tween 20), with each wash lasting 10 min with light shaking. Secondary antibody was incubated at a ratio of 1 μL per 10 mL of PBST-BSA for 1 h at room temperature with gentle shaking. Signal detection was achieved by adding Amersham electrochemiluminescent detection reagents (GE Healthcare, Buckinghamshire, UK) and imaging with the LAS 3000 camera system (Fujifilm). Western blot characterization of the initial IMAC-based purification using the MTBR-binding K9JA antibody followed a previously published protocol.⁵⁹

ThT Fluorescence Assay. Ten microliters each of heparin (MW 6000) and ThT (to yield 25 μM and 20 μM final concentrations, respectively) was added to 80 μL of tau K18 (9 μM final concentration) per well of a 96-well plate (Greiner Bio-One #655077). Aggregation was monitored for 15 h by measuring ThT fluorescence at 440 and 510 nm in a CLARIOstar microplate reader (BMG Lab Tech, Ortenberg, Germany). Sensitivity was set to 20% to prevent signal saturation. Reading using the top optic setting was obtained every 5 min after 200 rpm shaking for 30 s. The ThT signal at each time point was normalized to that of buffer only by subtracting the absorbance value of the blank at each identical time point from those of samples. Unless otherwise stated, the buffer used in all assays was 10 mM sterile-filtered sodium phosphate buffer pH 7.4.

Filament Assembly and Characterization by Negative-Stain TEM. Freshly purified samples (30 μM) were incubated with 63 μM heparin for 5 or 16 days at 37 °C without shaking. Samples were then diluted 1:2 with 10 mM sodium phosphate buffer pH 7.4 and analyzed by TEM as described.⁵⁹ The ImageJ freeware was used for filament length and width analysis as follows: after setting the scale, the freehand line tool was used to trace filament architecture, and the length was then measured. The straight line tool was used for width measurements. For filaments of apparently even width, measurements were taken once from random positions. For those coalescing from two filaments or of otherwise uneven widths, measurements were taken from representative spots as described in [Figure 9D](#). We followed the terminology used by Xu et al. (2010)⁷⁴ in characterizing fibril structures.

CD Spectroscopy. Tau K18 samples (200 μL) diluted in 10 mM sodium phosphate buffer pH 7.4 were transferred to a 1 mm path-length cuvette and analyzed on a Jasco J-815 CD spectropolarimeter (Jasco Inc., Maryland, USA) with 1 s response time, 0.1 nm data pitch, and 100 nm/min scan speed. Averaged spectral accumulations were collected at 190 to 280 nm wavelengths, with the high-tension voltage maintained at ≤ 550 V.

CD analysis of monomers involved 15 μM K18 WT and K18 C291A/C322A/I260C. For time-dependent CD traces, data were taken on 20 μM monomers on day 0 after which an 83 μM final concentration of heparin was added, and the mixture was incubated at 37 °C without shaking. CD measurements were repeated on days 4 and 8. In sedimentation assays, filament samples were ultracentrifuged at 100,000 $\times g$ for 1 h at 4 °C, the supernatant was removed, and the

pellet was resuspended in 200 μL of 10 mM sodium phosphate buffer pH 7.4 for CD analysis.

Statistics. Data are expressed as mean \pm standard error of the mean (SEM) from $n = 3$ experiments, unless otherwise stated. Data analysis and graph preparation were performed using Prism 7 (GraphPad Inc., California, USA). Statistical comparison of filament features was performed using the Mann–Whitney test at the 95% confidence interval, as the data did not follow Gaussian distribution.

■ ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at <https://pubs.acs.org/doi/10.1021/acscchemneuro.0c00469>.

Description of library of tau plasmids generated, primers used to generate plasmids for expressing and sequencing tau protein variants, Western blot and SEC characterization of expression and purification of tau variants, and CD spectroscopy characterization of soluble and insoluble tau aggregates ([PDF](#))

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Author Contributions

T.K.K., S.K., E.H., J.L.R., D.A.N., and B.B. performed the experiments and analyzed data. K.H., H.Z., K.B., E.J.H., and K.G.M. provided supervision, intellectual contributions, and contributed reagents. T.K.K. wrote the first manuscript draft which was edited by D.A.N., H.Z., K.B., E.J.H., and K.G.M. All authors read, substantially contributed to, and approved the manuscript for submission.

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Notes

The authors declare no competing financial interest.

Supporting research data regarding 44 sequence-verified plasmids for this article may be requested from the corresponding author by completing the following form: https://docs.google.com/forms/d/e/1FAIpQLSfcr1z3LF0v8EVxvPoyf6tKaWldktX_q-mPILPySEBfI99OMw/viewform?usp=pp_url.

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ABBREVIATIONS

AD, Alzheimer's disease; AFM, atomic force microscopy; CV, column volumes; FL, full length; FTD, frontotemporal dementia; IMAC, immobilized metal affinity purification; K18, four-repeat tau fragment; MRE, mean residue ellipticity; IPTG, isopropyl β -D-1-thiogalactopyranoside; MT, microtubule; MTBR, MT-binding repeat; NFT, neurofibrillary tangle; PCR, polymerase chain reaction; PHF, paired helical filament; RFU, relative fluorescent units; SF, straight filament; SEC, size exclusion chromatography; SEM, standard error of the mean; TEM, transmission electron microscopy; ThT,

thioflavin T; WT, wild type; WB, Western blot(ting); 4R, four-repeat; 3R, three-repeat

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